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2020-08

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Kiheri , H , Velmala , S , Pennanen , T , Timonen , S , Sietiö , O-M , Fritze , H , Heinonsalo , J , van Dijk , N , Dise , N & Larmola , T 2020 , ' Fungal colonization patterns and enzymatic activities of peatland ericaceous plants following long-term nutrient addition ' , Soil Biology & Biochemistry , vol. 147 , 107833 . <https://doi.org/10.1016/j.soilbio.2020.107833>

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<http://hdl.handle.net/10138/317317>

<https://doi.org/10.1016/j.soilbio.2020.107833>

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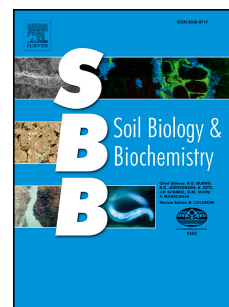
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Fungal colonization patterns and enzymatic activities of peatland ericaceous plants following long-term nutrient addition

Heikki Kiheri, Sannakajsa Velmala, Taina Pennanen, Sari Timonen, Outi-Maaria Sietiö, Hannu Fritze, Jussi Heinonsalo, Netty van Dijk, Nancy Dise, Tuula Larmola



PII: S0038-0717(20)30130-9

DOI: <https://doi.org/10.1016/j.soilbio.2020.107833>

Reference: SBB 107833

To appear in: *Soil Biology and Biochemistry*

Received Date: 4 November 2019

Revised Date: 17 April 2020

Accepted Date: 21 April 2020

Please cite this article as: Kiheri, H., Velmala, S., Pennanen, T., Timonen, S., Sietiö, O.-M., Fritze, H., Heinonsalo, J., van Dijk, N., Dise, N., Larmola, T., Fungal colonization patterns and enzymatic activities of peatland ericaceous plants following long-term nutrient addition, *Soil Biology and Biochemistry* (2020), doi: <https://doi.org/10.1016/j.soilbio.2020.107833>.

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1     **Fungal Colonization Patterns and Enzymatic Activities of Peatland**

2     **Ericaceous Plants Following Long-Term Nutrient Addition**

3

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**Abstract**

Northern peatlands are often dominated by ericaceous shrub species which rely on ericoid mycorrhizal fungi (ERM) for access to organic sources of nutrients, such as nitrogen (N) and phosphorus (P), and host abundant dark septate endophytes (DSE). Relationships between hosts and fungal symbionts may change during deposition of anthropogenic N and P. We studied the long-term effects of N and P addition on two ericaceous shrubs, *Calluna vulgaris* and *Erica tetralix*, at Whim Bog, Scotland by analyzing fungal colonization of roots, enzymatic activity, and fungal species composition. Unexpectedly, the frequency of typical ERM intracellular colonization did not change while the occurrence of ERM hyphae tended to increase and DSE hyphae to decrease. Our findings indicate that altered nutrient limitations shift root associated fungal colonization patterns as well as affecting ericaceous root enzyme activity and thereby decomposition potential. Reduction of recalcitrant fungal biomass in melanized DSE may have implications for peatland C sequestration under nutrient addition.

**Keywords:** ericoid mycorrhizae; peatland; nutrient limitation; enzyme activity; mycorrhizal colonization; dark septate endophyte; nitrogen deposition; *Sphagnum*

Declarations of interest: none

## 49    **1 Introduction**

50    Peatlands in the Northern hemisphere are often nutrient poor  
51    ecosystems characterized by acidic, anoxic, water saturated conditions  
52    with considerable nitrogen (N) and phosphorus (P) limitations (Aerts et  
53    al., 2001). These conditions and the accumulation of recalcitrant  
54    vegetation litter containing high concentrations of phenolic compounds  
55    and humic acids inhibitory to microorganisms and vegetation are  
56    considered to largely suppress decomposition (Leake and Read, 1990;  
57    Painter, 1991; Read et al., 2004; van Breemen, 1995). The challenging  
58    conditions in peatlands support a unique diversity of vegetation, with  
59    ericaceous species comprising one of the most dominant ground cover  
60    groups. Ericaceous shrubs are largely dependent on ericoid mycorrhizal  
61    fungi (ERM) to provide access to organic N and P which they provide in  
62    exchange for photosynthetic carbon (C) from the host plant (Smith and  
63    Read, 2008). The ERM fungi are capable of accessing organic N and P via  
64    a large variety of degradative enzymes which act primarily on plant cell  
65    wall components (Perotto et al., 2018), demonstrating a potential  
66    versatility more comparable to saprotrophs than to other types of  
67    mycorrhizae. These ericaceous species are also host to abundant dark  
68    septate endophytes (DSE) with extracellular enzyme capabilities  
69    potentially capable of improving host nutrient uptake (Mandyam and  
70    Jumpponen, 2005, 2014; Upson et al., 2009).

71

72 Over the past 150 years atmospheric deposition of N and P in forms  
73 easily accessible to plants has been increasing through combustion of  
74 fossil fuels and agricultural fertilization (Galloway et al., 2013; Galloway  
75 et al., 2003; Tipping et al., 2014; Wang et al., 2015). As nutrient  
76 limitations are alleviated, ericaceous reliance on ERM fungi may be  
77 reduced, potentially altering the symbiont community and leading to  
78 the loss of mycorrhizal symbionts. As mycorrhizal symbionts likely play a  
79 role in protection against pathogens (Vohník et al., 2016; Weiß et al.,  
80 2016), host species may in turn become more vulnerable to infection.  
81 Furthermore, the long-term effects of an altered nutrient balance in  
82 peatlands may include reduced nutrient acquisition competitiveness for  
83 ericaceous species. The ERM fungi are not as efficient decomposers as  
84 free-living saprotrophs, which may be naturally kept in check by  
85 nutrient limitations and direct competition with ERM fungi (Averill et al.,  
86 2014). Reduction of nutrient limitations may free the saprotrophic  
87 species' decomposition potential, leading to their dominance and the  
88 decline of ERM species and their host plants.

89

90 The selective pressure of N deposition on the symbiont community  
91 highlights a risk to peatland plant diversity, potentially leading to similar  
92 large-scale community shifts as seen in the continental level decline of  
93 ectomycorrhizal tree species and the increase in arbuscular tree species  
94 described by Averill et al. (2018). Any large-scale changes to peatland  
95 microbial and plant communities risk changing the status of peatlands

96 as net C sinks to net sources of greenhouse gas emissions (Andersen et  
97 al., 2013), which holds a globally significant potential when considering  
98 that peatlands sequester nearly one third of global soil organic carbon C  
99 (Gorham, 1991). The risk to C sink potential has been indicated by  
100 Larmola et al. (2013), who found that ecosystem C uptake did not  
101 increase in a long-term nutrient addition experiment at a nutrient-poor  
102 peatland in Canada simulating atmospheric N deposition.

103

104 The Whim Bog experimental site, located in the Scottish Borders, was  
105 established in 2001 to study the effects of different N forms and P  
106 addition on an ombrotrophic peat bog (Sheppard et al., 2004). This site  
107 allowed us to study nutrient addition effects on the ericaceous species  
108 *Calluna vulgaris* and *Erica tetralix* and their root associated fungi. The  
109 primary goals of this study were to characterize the frequency and  
110 morphology of mycorrhizal colonization in these ericaceous shrub  
111 species, assess ericoid mycorrhizal root enzyme capability related to  
112 organic matter degradation, and identify their fungal symbionts under  
113 changing nutrient availability. We hypothesized that: (1) the frequency  
114 of microscopically observed fungal colonization in ericaceous shrub  
115 roots is reduced across both forms of N and NP nutrient addition  
116 treatments, reflecting a reduction in reliance on symbionts for nutrient  
117 uptake; (2) nutrient addition treatments alter root associated fungal  
118 diversity, as determined by morphotypic analysis and ITS sequencing;

119 and (3) the activities of ericoid mycorrhizal root surface enzymes reflect  
120 treatment nutrient limitations.

121

## 122 **2 Materials and Methods**

### 123 **2.1 Study Site**

124 The study site, Whim Bog, located in the Scottish Borders, UK (Latitude:  
125 55.76670, Longitude: -3.26667), has undergone nutrient addition  
126 treatment since 2001. The study site, Whim Bog, located in the Scottish  
127 Borders, UK (Latitude: 55.76670, Longitude: -3.26667), has undergone  
128 nutrient addition treatment since 2001. The four nutrient addition  
129 treatments included in this study received annually  $6.4 \text{ g N m}^{-2}$  either as  
130 sodium nitrate ( $\text{NaNO}_3$ ) or as ammonium chloride ( $\text{NH}_4\text{Cl}$ ), both with  
131 and without P and K (as  $\text{K}_2\text{HPO}_4$ ).  $\text{K}_2\text{HPO}_4$  was added at a 1:14 P:N ratio  
132 to represent the ratio found in amino acids (See Sheppard et al. (2004)  
133 and Levy et al. (2019) for details). The ambient deposition in controls  
134 was  $0.8 \text{ g N m}^{-2}$ . Precipitation collected at the site was mixed with  
135 standard solutions to the required treatment concentrations. When  
136 adequate precipitation was collected an automated sprayer-system  
137 applied the treatment to the plots, simulating natural rainfall. Natural  
138 precipitation was not excluded from plots. Plots received 15 years of  
139 nutrient addition resulting in a  $96 \text{ g N m}^{-2}$  cumulative load.

140

### 141 **2.2 Site Measurements and Sampling**



142 In August 2016, plant species composition and abundance for each plot  
143 were measured via the point-intercept method using a 0.36 m<sup>2</sup> frame on  
144 permanent vegetation quadrats established on site. The frame was  
145 placed at a height of ca. 1 m relative to the surface of the plot and a  
146 graduated pin was used to measure the frame height and vertical  
147 location of each vegetation point for 61 intercepts as described in  
148 Larmola et al. (2013). Water table (WT) depth for each plot was  
149 measured from holes present after extraction of ingrowth cores used in  
150 a separate study, relative to moss surface height.  
151  
152 Triplicate plots per treatment and controls were sampled in November  
153 2016 for *Calluna vulgaris* (L.) Hull and *Erica tetralix* (L.) plant roots from  
154 one individual plant per plot. Fine root sections were collected from  
155 several points throughout the root system of each sampled plant and  
156 stored at 8 °C prior to transport, followed by storage at -20 °C. Each root  
157 sample was split into two subsamples: one for microscopy and the other  
158 for enzymatic and subsequent molecular analyses.  
159  
160 Surface peat from triplicate plots per treatment and controls was  
161 sampled to a depth of 20 cm in each plot and stored at 8 °C prior to  
162 transport, followed by storage at -20 °C. After melting, the pH of each  
163 sample was measured following homogenization with deionized water  
164 at a 1:4 ratio.  
165

### 2.3 Microscopy

The mycorrhizal status of the ericaceous shrub species *C. vulgaris* and *E. tetralix* was determined via light microscopy and Trypan Blue staining as described in Kiheri et al. (2017). Using the magnified intersections method described by McGonigle et al. (1990), with one slide per species from each of three replicate plots per treatment and controls (15 samples per species), using 300 counts per slide, mycorrhizal colonization was quantified according to different morphological categories, described in Table 1. These categories were used to estimate differences between the frequencies of potential ericoid mycorrhizal (ERM) hyphae and typical dark septate endophyte (DSE) hyphae. In our study, only those fungi which were robust in structure, heavily melanized and displaying septa were assessed to be DSE. This was necessary to prevent misclassification of those ERM species which produce melanized and septate hyphae (Vohník and Albrechtová, 2011).

Table 1. Microscopic morphological categories of ericaceous root associated fungi.

Hyphal Type	Quantification Category	Morphological Characteristics
	ERM0/DSE0	No fungal presence
<b>Potential ericoid mycorrhizae (ERM)</b>	ERM1	Intracellular coiling
	ERM2	Intracellular hyphae
	ERM3	Colonizing surface hyphae
	ERM4	Extracellular surface hyphae
<b>Dark Septate Endophyte (DSE)</b>	DSE1	Intracellular coiling
	DSE2	Intracellular hyphae
	DSE3	Colonizing surface hyphae
	DSE4	Extracellular surface hyphae

184

185 Hereon, putative ericoid mycorrhizal morphotypes are referred to with  
186 categories ERM1-4 and dark septate endophytes with DSE1-4 (Table 1).  
187 Of the morphotypic categories, only ERM category 1 (intracellular  
188 coiling) was interpreted as mycorrhizal colonization frequency.  
189 Categories 2-4 were interpreted as potential changes in fungal diversity  
190 and function.

191

#### 192 2.4 Enzyme Assays

193 The enzymatic activities of root samples from *C. vulgaris* and *E. tetralix*  
194 were determined using a multi-enzyme assay described by Pritsch et al.  
195 (2011), originally performed for assessing ectomycorrhizal exo-enzyme  
196 potential. From triplicate plots of each of the five treatments, rhizomes  
197 of both ericaceous species were each sampled for 9 individual ca. 1 cm  
198 root pieces, for a total of 270 root samples ( $n = 5$  treatments  $\times$  3 plots  $\times$   
199 2 species  $\times$  9 root samples). To measure eight different hydrolytic and  
200 oxidative root surface enzyme potentials we used the method described  
201 by Velmala et al. (2014) for fluorescences representing the potential  
202 activities of leucine aminopeptidase (EC 3.4.11.1), hemicellulases via  $\beta$ -  
203 glucuronidase (EC 3.2.1.31) and  $\beta$ -xylosidase (EC 3.2.1.37), cellulases via  
204 cellobiohydrolase (EC 3.2.1.91) and  $\beta$ -glucosidase (EC 3.2.1.21),  
205 chitinase via N-acetylglucosaminidase (EC 3.2.1.14), and acid  
206 phosphatase (EC 3.1.3.2). Samples were incubated at room temperature  
207 in the dark and under agitation at 180 rpm on a tabletop shaker, with

208 each root piece in individual wells of 96-well filter plates (30–40  $\mu\text{m}$   
209 mesh size, AcroPrep™ 96 Filter Plate; PALL, Port Washington, NY, USA)  
210 in buffers containing enzyme specific 7-amino-4-methylcoumarine  
211 (AMC) or 4-methylumbelliferone (MU) substrates. Incubation times for  
212 each enzyme followed the protocol of Pritsch et al. (2011). The  
213 respective substrates used were Leucine-AMC, MU-xylopyranoside, MU-  
214  $\beta$ -D-glucuronide, MU-cellobiohydrofuran, MU-N-acetyl- $\beta$ -D-  
215 glucosaminide, MU- $\beta$ -D-glucopyranoside, and MU-phosphate. After  
216 incubation, substrate solutions were collected by centrifugation with a  
217 96-well plate adapter at 3200 rpm onto Optiplate-96F reading plates  
218 (Perkin-Elmer, Waltham, MA, USA) containing stop buffer (pH 10). Each  
219 substrate's fluorescence was measured using a Victor<sup>3</sup> 1420 multilabel  
220 plate counter (Perkin-Elmer, Waltham, MA, USA) at an excitation  
221 wavelength of 355 nm and an emission wavelength of 460 nm. Standard  
222 solutions were prepared using aminomethylcoumarin (AMC) and 4-  
223 methylumbelliferone (MUF) and used to calculate enzymatic activities  
224 from concentrations of released AMC or MUF according to their  
225 respective substrates. All standard and enzyme substrate solutions were  
226 purchased from Sigma-Aldrich (Merck KGaA, Darmstadt, Germany).

227

228 Further, laccase (EC 1.10.3.2) activity was used as an indicator of lignin  
229 modification activity and was determined by incubation in buffer  
230 containing diammonium 2,2'-azinobis-3-ethylbenzothiazoline-6-  
231 sulfonate (ABTS) and colorimetric measurement using a Tecan Infinite

232 M200 PRO Multimode Reader (Tecan Trading AG, Männedorf,  
233 Switzerland).

234

235 Following enzymatic measurements, the individual root pieces were  
236 scanned at high resolution (650 dpi) and their surface area measured  
237 using winRHIZO Pro (ver. 2017, Regent Instruments Inc.) software.  
238 These surface area values were used to convert enzymatic activity to  
239  $\text{pmol mm}^{-2} \text{ min}^{-1}$ .

240

## 241 2.5 Molecular Methods and Sequencing

242 From triplicate plots of each of the five treatments, rhizomes of both  
243 ericaceous species were each sampled for 9 individual ca. 1 cm root  
244 pieces, for a total of 270 root samples ( $n = 5 \text{ treatments} \times 3 \text{ plots} \times 2$   
245  $\text{species} \times 9 \text{ root samples}$ ). Following enzyme assays and winRHIZO  
246 analysis on these samples, a randomized subset of 1 of 3 treatment  
247 replicates was taken from the 270 samples for sequencing. These 90  
248 samples, plus four repeated samples, were then directly amplified using  
249 a Phire Plant Direct PCR kit's plant leaf protocol (Thermo Fisher  
250 Scientific, Waltham, MA, USA) and PCR using the ITS1F (Gardes and  
251 Bruns, 1993) and ITS4 (White et al., 1990) primer pair. Individual root  
252 pieces were manually crushed using sterile pestles in provided dilution  
253 buffer, in order to release fungal cells both on the root surface and  
254 within the root structure, and 1  $\mu\text{l}$  of each mixture was then used as a  
255 template in a 20  $\mu\text{l}$  PCR reaction. Cycling conditions for the Direct PCR

256 were: initial denaturation 98 °C 5 min, 40 cycles (98 °C for 5 sec, 57 °C  
257 for 5 sec, 72 °C for 20 sec), and final extension 72 °C for 1 min. The  
258 Direct PCR products were separated in a 2% agarose gel in 1X TAE buffer  
259 at 120V for 2 hours and each ITS band excised and purified using a  
260 Nucleospin® Gel and PCR Clean-Up kit (Macherey-Nagel GmbH & Co.  
261 KG, Düren, Germany). The purified products were then further amplified  
262 using a DreamTaq PCR mastermix (Thermo Fisher Scientific, Waltham,  
263 MA, USA) by using 1 ul of Direct PCR product as the template in a 20 µl  
264 DreamTaq PCR reaction with the same ITS1F-ITS4 primer pair. The  
265 DreamTaq PCR was performed with the following program: initial  
266 denaturation 95 °C for 3 min, 35 cycles (95 °C for 30 sec, 57 °C for 30  
267 sec, 72 °C for 1 min), and final extension 72 °C for 10 min. This was done  
268 as a quick method to identify common fungal species in each root  
269 sample which could then be linked to their associated root surface  
270 enzyme activities. These ITS products were Sanger sequenced using the  
271 ITS4 primer (Macrogen Europe, Amsterdam, NL). This produced ITS  
272 fragments ranging from 98 to 884 bp in length. This method resulted in  
273 the successful sequencing of 60% of (57 out of 94) ITS products  
274 (Supplementary File 1), of which 38 sequences were more than 200 bp  
275 in length and of high enough quality required for NCBI Gen Bank  
276 submission. The sequences are disposed to Gen Bank under the  
277 accession numbers **MN059889-MN059927**. See Supplementary File 1  
278 for all 57 FASTA sequences. The high proportion of sequencing failure  
279 was likely due to mixed ITS products from multiple fungal species and

280 inhibitory chemistry. Sequence identification was performed using the  
281 Unite massBLASTer analysis (Nilsson et al., 2019) on manually trimmed  
282 sequences. Analyzed sequences had 86-100% sequence similarity to  
283 existing reference or representative sequences within the INSD or  
284 Environmental databases and the most likely species hypotheses (SH)  
285 were selected as their fungal identities. Functional roles were then  
286 assigned to these fungal identities according to the web based FUNGuild  
287 bioinformatic tool (Nguyen et al., 2016).

288

## 289 2.6 Statistical Analyses

290 Hereon,  $\alpha$  level for statistical significance is defined as  $p \leq 0.05$  and  
291 indicative as  $0.05 < p \leq 0.1$  in all cases, and the term significant  
292 specifically indicates statistical significance as  $p \leq 0.05$ . All analyses were  
293 calculated using values from treatment means ( $n=3$ ). Differences  
294 between treatments, for each host plant separately, were determined  
295 by analysis of variance (ANOVA) on logarithmically transformed data  
296 followed by pairwise comparisons using the parametric Tukey's HSD and  
297 nonparametric Games-Howell post hoc tests, using IBM SPSS Statistics  
298 25. Fungal-Enzyme activity profiles in Figure 8 were prepared using  
299 OriginPro 2018 by assigning fungal identifications to enzyme activities  
300 on an individual root piece basis.

301

302 The combined data for both host plants' mycorrhizal morphotype  
303 categories ERM1-4 and DSE1-4 (Table 1), root enzyme activities, plot

level vegetation abundance, surface peat pH, and water table depth were analyzed using correlation analyses. Correlation analyses were performed using rcorr-function with Spearman rank based correlation from package Hmisc v4.1-1 (Harrell et al., 2014) and plotted using corrplot-function from package corrplot v0.84 (Wei and Simko, 2016) in the R programming environment (R Core Team, 2017).

### 3 Results

#### 3.1 Vegetation, Peat pH, and Water Table

In all four nutrient addition treatments, the dominant ericaceous shrub, *Calluna vulgaris*, tended to decrease in abundance while *Erica tetralix* tended to increase, especially in NaNO<sub>3</sub> treatments (Table 2). Nutrient addition treatments showed decreasing trends in the abundance of *Sphagnum*, when compared with controls. The sedge *Eriophorum vaginatum* showed increasing trends in abundance in all treatments except NaNO<sub>3</sub>, when compared with controls. The high abundance of reported *E. vaginatum* in the NaNO<sub>3</sub>+PK treatment was largely an effect of one plot where the point-intercept measurements were performed within a large *E. vaginatum* tussock. Furthermore, the different forms of N addition were found to cause opposite changes in peat pH, with NaNO<sub>3</sub> increasing pH by ca. 0.2-0.3 units and NH<sub>4</sub>Cl decreasing pH by ca. 0.2-0.4 units (Table 2). Effects of treatments on peat pH were found to be statistically significant ( $F_{(4,10)}=6.410$ ), with pairwise comparisons finding statistically significant differences between NaNO<sub>3</sub> and NH<sub>4</sub>Cl+PK



treatments and between  $\text{NaNO}_3$ +PK and  $\text{NH}_4\text{Cl}$ +PK treatments. Mean water table (WT) depth, measured relative to moss surface, was eight cm closer to the moss surface in  $\text{NaNO}_3$ +PK treatments than in control plots, while  $\text{NH}_4\text{Cl}$  treatments affected WT the least. This was likely affected by the loss of moss abundance and subsequent subsidence in  $\text{NaNO}_3$ +PK treatments.

334

Table 2. Vegetation abundance (hits per  $\text{m}^2$ ), surface (0-20cm) peat pH, and treatment water table (WT) depth with  $\pm 1$  standard deviation,  $n=3$ . Different superscript letters indicate significant differences ( $P<0.05$ ) compared with the other treatments. Statistically significant differences were only observed for pH values ( $F_{(4,10)}=6.410$ ).

	<i>Calluna vulgaris</i>	<i>Erica tetralix</i>	<i>Eriophorum vaginatum</i>	Other Vasc. spp.	<i>Sphagnum</i>	Other Mosses	pH	WT (cm)
<b>Control</b>	212.6 $\pm$ 33.6	4.0 $\pm$ 4.9	42.3 $\pm$ 32.0	2.0 $\pm$ 2.1	24.3 $\pm$ 17.5	28.3 $\pm$ 16.7	4.1 $\pm$ 0.0 <sup>ab</sup>	16.0 $\pm$ 12.4
<b><math>\text{NaNO}_3</math></b>	156.6 $\pm$ 72.3	57.0 $\pm$ 28.7	23.3 $\pm$ 2.3	5.6 $\pm$ 7.3	11.6 $\pm$ 16.4	25.3 $\pm$ 1.6	<b>4.4 <math>\pm</math> 0.2<sup>a</sup></b>	12.6 $\pm$ 9.8
<b><math>\text{NaNO}_3</math> +PK</b>	61.0 $\pm$ 41.4	11.0 $\pm$ 14.8	205.3 $\pm$ 182.1	5.0 $\pm$ 2.9	0.0	35.0 $\pm$ 11.3	<b>4.3 <math>\pm</math> 0.1<sup>a</sup></b>	8.0 $\pm$ 1.4
<b><math>\text{NH}_4\text{Cl}</math></b>	221.6 $\pm$ 49.7	19.0 $\pm$ 12.3	57.0 $\pm$ 26.1	0.0	10.6 $\pm$ 5.2	20.3 $\pm$ 19.7	3.9 $\pm$ 0.1 <sup>ab</sup>	16.0 $\pm$ 2.6
<b><math>\text{NH}_4\text{Cl}</math> +PK</b>	163.3 $\pm$ 72.6	24.6 $\pm$ 23.7	113.3 $\pm$ 64.5	1.3 $\pm$ 1.8	9.6 $\pm$ 8.01	24.6 $\pm$ 8.1	<b>3.7 <math>\pm</math> 0.1<sup>b</sup></b>	24.0 $\pm$ 4.2

340

### 3.2 Root Associated Fungal Morphology and Frequency

The mean proportion of root intersects which were microscopically quantified as presenting fungal colonization increased from 78% and 83% in control plots for *C. vulgaris* and *E. tetralix* roots, respectively, to 89% and 91% in nutrient addition plots, respectively. Of these root intersections which contained fungal structures, both ericaceous host

347 species showed trends of increasing ERM hyphal frequency and  
 348 decreasing DSE hyphal frequency when under nutrient addition (Figure  
 349 1). Nutrient addition increased mean ERM hyphal frequency in *C.*  
 350 *vulgaris* and *E. tetralix* roots by 31% and 16%, respectively. However,  
 351 mean DSE hyphal frequency decreased by 30 % in *C. vulgaris* roots and  
 352 22% in *E. tetralix* roots. Though these trends are considerable, they  
 353 were not found to be statistically significant due to high natural  
 354 variation between samples.  
 355  
 356 Intracellular hyphal frequency (ERM2, Table 1) of *C. vulgaris* roots  
 357 significantly ( $F_{(4,10)}=11.406$ ) increased by ca. two-fold under nutrient  
 358 addition when comparing controls to  $\text{NaNO}_3$ +PK,  $\text{NH}_4\text{Cl}$ , and  $\text{NH}_4\text{Cl}$ +PK,  
 359 as well as increasing by approximately half under  $\text{NaNO}_3$  addition, which  
 360 was a statistically indicative change (Figure 2). *C. vulgaris* roots showed  
 361 no statistically significant increases in cells containing typical ERM  
 362 intracellular coiling (ERM1), root surface colonizing hyphae (ERM3), or  
 363 extracellular surface hyphae (ERM4) under nutrient addition. *C. vulgaris*  
 364 DSE extracellular hyphal frequency (DSE4) decreased significantly under  
 365  $\text{NH}_4\text{Cl}$  addition to less than 10% that of controls ( $F_{(4,10)}=4.654$ ) (Figure 3).  
 366 The same analysis for *E. tetralix* roots showed similar trends of  
 367 increasing ERM morphotype frequencies under nutrient addition (Figure  
 368 2), while DSE morphotype frequencies decreased (Figure 3).  
 369 Interestingly, a statistically indicative positive correlation was found

370 between *E. tetralix* root surface DSE colonization frequency (DSE3) and  
371 plot *Sphagnum* abundance ( $r = 0.58$ ) (Figure 5).

372

### 373 3.3 Root Enzymatic Activity

374 The different forms of N addition,  $\text{NH}_4^+$  and  $\text{NO}_3^-$ , induced variable and  
375 opposing effects on C and N acquiring enzymes on the two ericaceous  
376 shrubs' root surfaces, when compared to controls (Figures 6&7).

377 Treatment with  $\text{NaNO}_3$  tended to reduce all *C. vulgaris* root enzymatic  
378 activities except acid phosphatase, suggesting that *C. vulgaris* or its root  
379 associated fungi are more sensitive to  $\text{NaNO}_3$  than *E. tetralix*, which did  
380 not show this effect. In *C. vulgaris* roots under  $\text{NH}_4\text{Cl}$  addition, the  
381 activities of the C acquiring enzymes  $\beta$ -xylosidase and  $\beta$ -glucosidase did  
382 not change compared to controls, while activity of  $\beta$ -glucuronidase  
383 tended to increase two-fold and N-acetylglucosaminidase tended to  
384 decrease by nearly half. In contrast, the effects of both forms of N on *E.*  
385 *tetralix* roots were similar for all C and N acquiring enzymes, while  
386 leucine aminopeptidase activity tended to be suppressed under  $\text{NH}_4\text{Cl}$   
387 addition to less than 25% of controls and N-acetylglucosaminidase  
388 decreased by less than half, compared to controls. Laccase activity was  
389 not detected in any samples.

390

391 Compared to control plots, addition of both forms of N alone tended to  
392 induce an approximately 25% increase in acid phosphatase activity in  
393 both plant species (Figures 6&7). In contrast, treatments with additional

394 PK reduced acid phosphatase activity to  $1/3^{\text{rd}}$  of controls in both  
 395 ericaceous species (Figure 7). In *C. vulgaris* roots, the effect of nutrient  
 396 additions on acid phosphatase activity was statistically significant  
 397 ( $F_{(4,10)}=8.163$ ), with significant differences between  $\text{NaNO}_3$  and  
 398  $\text{NaNO}_3+\text{PK}$  and  $\text{NH}_4\text{Cl}+\text{PK}$  treatments, as well as between  $\text{NH}_4\text{Cl}$  and  
 399  $\text{NaNO}_3+\text{PK}$  and  $\text{NH}_4\text{Cl}+\text{PK}$  treatments. In *E. tetralix* roots the effect of  
 400 nutrient addition on acid phosphatase activity was significant  
 401 ( $F_{(4,10)}=11.400$ ), with significant differences between control and  
 402  $\text{NH}_4\text{Cl}+\text{PK}$  treatments, between  $\text{NaNO}_3$  and  $\text{NaNO}_3+\text{PK}$  and  $\text{NH}_4\text{Cl}+\text{PK}$   
 403 treatments, and also between  $\text{NH}_4\text{Cl}$  and  $\text{NaNO}_3+\text{PK}$  and  $\text{NH}_4\text{Cl}+\text{PK}$   
 404 treatments.  
 405  
 406 Interestingly, in both ericaceous species the NPK treatments induced  
 407 higher, although statistically non-significant, activities in many C and N  
 408 acquiring enzymes compared to N alone (Figures 6&7). The exceptions  
 409 to this were the suppression of  $\beta$ -xylosidase and  $\beta$ -glucuronidase  
 410 activities in *C. vulgaris* roots under  $\text{NH}_4\text{Cl}+\text{PK}$  addition. Additionally,  
 411 there was a significant positive correlation between *C. vulgaris* ERM  
 412 intracellular hyphal frequency (ERM2) and  $\beta$ -glucuronidase activity ( $r=$   
 413  $0.66$ ) and a statistically indicative positive correlation with  $\beta$ -glucosidase  
 414 activity ( $r= 0.50$ ) (Figure 4).  
 415  
 416 3.4 Ericaceous Root Associated Fungi

417 Sanger sequencing of Direct PCR ITS amplicons from individual root  
418 samples from both ericaceous species revealed several confirmed and  
419 putative ERM/DSE fungal species, as well as a range of possible  
420 endophytes, saprotrophs, and pathogens (Table 3). Fungal sequence  
421 identifications based on Species Hypotheses (SH) ranged from 86% to  
422 100% matches with reference sequences. Identifications of ascomycete  
423 fungi likely inhabiting these ericaceous roots as ERM symbionts included  
424 *Hyaloscypha hepaticicola* and *Hyaloscypha* sp., (names updated from  
425 *Rhizoscyphus ericae* and *Meliniomyces* sp., respectively, according to  
426 Fehrer et al. (2019)). Other members of the Leotiomycetes were also  
427 identified, including unidentified Helotiales, unidentified  
428 *Hyaloscyphaceae*, *Phialocephala sphaeroides*, and *Pseudogymnoascus*  
429 sp. Furthermore, identified basidiomycetes which are capable of the  
430 ERM lifestyle included members of the family *Serendipitaceae* (syn.  
431 clade B Sebacinales) and its member genus *Serendipita*. Interestingly,  
432 members of the Helotiales were found only in roots of both hosts from  
433 control,  $\text{NH}_4\text{Cl}$  or  $\text{NH}_4\text{Cl}+\text{PK}$  treatments while unidentified Pezizales  
434 members were only detected in  $\text{NaNO}_3+\text{PK}$  treatments. Other  
435 ericaceous root associated fungi which were possibly living saprotrophic  
436 or pathotrophic lifestyles are presented in Table 3.

Table 3. Fungal identifications from Sanger sequencing of Direct PCR ITS amplicons from *C. vulgaris* and *E. tetralix* roots. % ID values indicate range of similarity with reference sequences according to Unite database. Function assignment according to FUNGuild analysis. Sequences are from 38 samples with ITS amplicons >200 bp as required by Unite Gen Bank, see Supplementary File 1 for all 57 FASTA sequences. Sources of sequences listed by treatments and host species indicated with Cv and Et, numbers in brackets indicate the number of replicate plots with the same sequence, subscripts c=Control, 1=NaNO<sub>3</sub>, 2=NaNO<sub>3</sub>+PK, 3=NH<sub>4</sub>Cl, 4=NH<sub>4</sub>Cl+PK.

Phylum	Class	Order	Family	Species	% ID	Function	Reference Seq(s) SH	Source
Ascomycota	Archaeorhizomycetes	Archaeorhizomycetales	Archaeorhizomycetaceae	<i>Archaeorhizomyces</i> sp.	97.58	Sapro	KT768305 SH180923.07FU	Cv <sub>c</sub>
	Dothideomycetes	Capnodiales	<i>Mycosphaerellaceae</i>	<i>Acrodontium crateriforme</i>	95.16	Patho-Sapro	KX287271 SH214154.07FU	Cv <sub>4</sub>
			<i>Cladosporiaceae</i>	<i>Cladosporium</i> sp.	99.77	Patho-Sapro-Symbio	KX459429 SH212842.07FU	Cv <sub>1</sub>
	Leotiomycetes	Helotiales	<i>Hyaloscyphaceae</i>	<i>Hyaloscypha hepaticicola</i> *	99.07	Patho-Sapro-Symbio	FM172802 SH181107.07FU	Cv <sub>3</sub>
				<i>Hyaloscypha</i> sp.*	99.18-99.59	Sapro-Symbio	FM997935 SH025067.07FU DQ309217 SH214267.07FU	Cv <sub>c</sub> ; Cv/Et <sub>3</sub> ;Cv <sub>4</sub>
				Unidentified	99.79	Sapro	HF947840 SH004619.07FU	Et <sub>c</sub>
			<i>Vibriseaceae</i>	<i>Phialocephala sphaeroides</i>	91.06	Symbio	KC480051 SH204990.07FU	Et <sub>3</sub>
			Unidentified	Unidentified	88.89-99.61	-	HF947859 SH218310.07FU AF252840 SH211416.07FU HF947861 SH197071.07FU AY627806 SH201639.07FU	Cv <sub>c</sub> ; Cv/Et(2) <sub>3</sub>
							AF149078 SH183994.07FU DQ309240 SH143881.07FU	
		Rhytismatales	Unidentified	Unidentified	92.00-95.71	-		Cv/Et <sub>c</sub> ;Cv <sub>1</sub>
		Thelebolales	<i>Pseudeurotiaceae</i>	<i>Pseudogymnoascus</i> sp.	93.89	Patho-Sapro-Symbio	KP902680 SH183329.07FU	Cv <sub>2</sub>
	Pezizomycetes	Pezizales	Unidentified	Unidentified	95.94-100	-	JQ347011 SH203769.07FU	Cv(2)/ Et(2) <sub>2</sub>
	Sordariomycetes	Microascales	<i>Halosphaeriaceae</i>	Unidentified	94.64	Sapro	FJ524322 SH211311.07FU	Et <sub>2</sub>
	Taphrinomycetes	Taphrinales	<i>Taphrinaceae</i>	<i>Taphrina tormentillae</i>	99.14	Patho	KX516468 SH200748.07FU	Et <sub>c</sub>
	incertae sedis	incertae sedis	incertae sedis	<i>Xenochalara</i> sp.	96.04	Sapro	HM230882 SH202721.07FU	Et <sub>4</sub>
Basidiomycota	Agaricomycetes	Agaricales	<i>Entolomataceae</i>	<i>Entoloma cetratum</i>	97.14	Patho-Sapro-Symbio	KC898450 SH185814.07FU	Cv <sub>2</sub>
			<i>Tricholomataceae</i>	Unidentified	93.82	Patho-Symbio	KY701558 NA	Cv <sub>3</sub>
			<i>Strophariaceae</i>	<i>Pholiota</i> sp.	86.47	Sapro	HQ533029 SH219745.07FU	Et <sub>2</sub>
		Cantharellales	<i>Ceratobasidiaceae</i>	<i>Ceratobasidium</i> sp.	99.49	Patho-Sapro-Symbio	JN569114 SH220624.07FU	Cv <sub>3</sub>
		Jaapiales	<i>Vibriseaceae</i>	<i>Jaapia ochroleuca</i>	99.31	Sapro	UDB031153 SH190037.07FU	Et <sub>3</sub>
		Sebacinales	<i>Serendipitaceae</i> (Clade B Sebacinales)	<i>Serendipita</i> sp.	98.25-100	Symbio	GQ907110 SH003898.07FU HF947895 SH201953.07FU DQ309211 SH179088.07FU DQ309149 SH180008.07FU	Et <sub>c</sub> ; Cv(2) <sub>1</sub> ; Et(2) <sub>3</sub>
							HF947869 HF947915/DQ309208 SH179085.07FU	
			<i>Sebacinaceae</i>	Unidentified	99.66	Symbio	HQ154421 SH199330.07FU	Cv <sub>c</sub> ;Et <sub>2</sub>
		Thelephorales	<i>Thelephoraceae</i>	<i>Thelephora terrestris</i>	98.72	Symbio	KX438350 SH184510.07FU	Et <sub>3</sub>
Mortierellomycota	Mortierellomycetes	Mortierellales	<i>Mortierellaceae</i>	<i>Mortierella humilis</i>	100	Sapro-Symbio	KM504403 SH196779.07FU	Cv/Et <sub>3</sub>
				<i>Mortierella parvispora</i>	99.83	Sapro-Symbio	FN565295 SH193938.07FU	Et <sub>4</sub>
Rozellomycota	unidentified	Unidentified	Unidentified	Unidentified	98.23	-	KF297176 SH204524.07FU	Et <sub>3</sub>

\*Sequences identified as *Rhizoscyphus ericae* and *Meliniomyces* sp. updated to as *Hyaloscyphus hepaticicola* and *Hyaloscyphus* sp., respectively, according to Fehrer et al. (2019)

### 443 3.5 Ericaceous Root Enzyme Activity Profiles

444 When fungal sequence identifications were linked to the enzymatic  
 445 activities of their respective root samples, potential species related  
 446 enzyme activity patterns emerged (Table 3, Figure 8). *C. vulgaris* root  
 447 samples from NaNO<sub>3</sub>, NaNO<sub>3</sub>+PK, and NH<sub>4</sub>Cl+PK treatments and hosting  
 448 *Serendipitaceae* (Clade-B Sebaciniales) or *Serendipita* sp. produced the  
 449 highest detected leucine aminopeptidase activities. A *C. vulgaris* root  
 450 sample in the NH<sub>4</sub>Cl+PK treatment and hosting *Hyaloscypha* sp. was  
 451 highly active in β-xylosidase, N-acetylglucosaminidase, and β-  
 452 glucosidase, while samples hosting *Hyaloscypha* sp. in control and NH<sub>4</sub>Cl  
 453 treatments were ca. 50-75 % less active for the same enzymes.  
 454 Additionally, a *C. vulgaris* root sample in the NH<sub>4</sub>Cl treatment  
 455 simultaneously hosting *Hyaloscypha hepaticicola* and *Ceratobasidium*  
 456 sp. was one of the most active samples, across all enzymes.  
 457  
 458 Interestingly, a root sample from *C. vulgaris* in the NaNO<sub>3</sub>+PK treatment  
 459 was linked to a fungal identification of *Pseudogymnoascus* sp., which  
 460 produced some of the highest activities for all enzymes in this  
 461 treatment, except leucine aminopeptidase. A sample of *C. vulgaris* root  
 462 from the NH<sub>4</sub>Cl+PK treatment which hosted the species *Acrodontium*  
 463 *crateriforme* indicated activities of cellobiohydrolase, N-  
 464 acetylglucosaminidase, and β-glucosidase several times higher than any  
 465 other samples measured. In the control treatment, *C. vulgaris* root  
 466 samples hosting a member of the Rhytismatales showed the highest

activity for leucine aminopeptidase and acid phosphatase while *E. tetralix* samples hosting Rhytismatales showed the highest activities for all enzymes in that treatment, except leucine aminopeptidase.

#### 4 Discussion

The increases in overall fungal colonization for both *Calluna vulgaris* and *Erica tetralix* under nutrient addition were unexpected and the unchanged frequency of ERM intracellular coiling showed that long-term N and NPK addition did not reduce mycorrhizal colonization. This suggests that the ericaceous host plants are unable to restrict fungal colonization of their roots, despite access to excess inorganic N and P. Alternatively, unchanged mycorrhizal colonization rates may indicate that the fungi provide benefits to the plant beyond N and P.

The observed reduction of *Sphagnum* abundance and the decreased frequency of DSE associated with ericaceous roots may be linked. DSE fungi have been shown to be a common occurrence in submerged aquatic plants (Kohout et al., 2012) and are capable of propagating and existing as saprobes of moss gametophytes (Day and Currah, 2011). This capability and their prevalence in aquatic plants suggest that DSE fungi are well adapted to periodic waterlogged conditions in peatlands, perhaps providing a niche during the annual senescence of their ericaceous hosts. A similar relationship between ERM fungi and liverworts, common species in peatlands, may also provide a niche



491 during host senescence (Kowal et al., 2018, 2015). As the loss of  
492 *Sphagnum* species is one of the most obvious effects of nutrient  
493 addition treatments (Bubier et al., 2007; Levy et al., 2019), this may  
494 remove an important ecosystem niche for DSE fungi. Our finding of  
495 positive correlation between *E. tetralix* root surface DSE colonization  
496 frequency and *Sphagnum* abundance supports this possibility.  
497  
498 The significantly increased frequencies of ERM hyphal morphotypes and  
499 decreased DSE hyphal morphotypes in ericaceous roots strongly  
500 suggests that long-term nutrient addition resulted in an altered fungal  
501 community. Furthermore, the different forms of N addition may have  
502 had selective effects on the fungal community as we identified different  
503 root associated Helotiales members only in controls,  $\text{NH}_4\text{Cl}$  or  $\text{NH}_4\text{Cl}+\text{PK}$   
504 treatments and members of the Pezizales only in  $\text{NaNO}_3+\text{PK}$  treatments.  
505 The Helotiales may prefer  $\text{NH}_4^+$  as a substrate compared to  $\text{NO}_3^-$ , as  
506 experimental evidence has shown for *Hyaloscypha hepaticicola* (Cairney  
507 et al., 2000). The presence of the Pezizales in only  $\text{NaNO}_3+\text{PK}$   
508 treatments may imply they prefer this N source, though their  
509 mycorrhizal status is unclear it has been suggested for some families by  
510 Hobbie et al. (2001).  
511  
512 Sequence identification of ITS amplicons from root samples of both  
513 ericaceous species revealed several ERM and DSE species commonly  
514 found to associate with ericaceous roots, as well as common peatland

515 saprotrophs and pathogens (Sietiö et al., 2018; Thormann, 2006;  
 516 Thormann and Rice, 2007). Both ericaceous species shared several  
 517 family and genus level groups, indicating a common symbiont  
 518 community among the ericaceous hosts. This is in line with the findings  
 519 of Kjølner et al. (2010) who showed that several ericaceous species in a  
 520 subarctic mire shared fungal communities when in close proximity.  
 521 While ERM symbionts are generally ascomycetes, recent studies have  
 522 found that the basidiomycete fungi *Serendipitaceae* (Clade-B  
 523 Sebaciniales) are common in ericaceous roots and capable of forming  
 524 mycorrhizal structures (Brundrett and Tedersoo, 2018; Vohník et al.,  
 525 2016; Weiß et al., 2016), and potentially capable of utilizing  
 526 photosynthetic C (Sietiö et al., 2018). Our identifications of members of  
 527 the *Serendipitaceae* and *Serendipita* sp. in both ericaceous hosts  
 528 support their likely role as root symbionts. Interestingly, in *C. vulgaris*  
 529 roots from the NaNO<sub>3</sub>+PK treatment we found a potential ERM  
 530 symbiont, *Pseudogymnoascus* sp., which is a genus that may form ERM  
 531 associations, as shown between *Pseudogymnoascus roseus* and  
 532 *Vaccinium angustifolium* (Dalpé, 1989).  
 533  
 534 Our findings on the varying effects of N addition on root surface  
 535 enzymatic activities in the two ericaceous species, *C. vulgaris* and *E.*  
 536 *tetralix*, indicate that they may have functionally different root  
 537 symbionts and decomposition potentials. In NPK treatments both  
 538 ericaceous plants displayed highly suppressed acid phosphatase activity

539 as the roots and root associated fungi did not need to access organic P  
540 sources. Conversely, N treatments increased acid phosphatase activity,  
541 reflecting the colimitation of N and P found in peatlands (Pinsonneault  
542 et al., 2016; Wang et al., 2015) and confirming that ericoid mycorrhizal  
543 root enzymatic activities reflect nutrient limitations. Almost every  
544 enzyme activity increased with additional P compared to either form of  
545 N alone, while  $\text{NaNO}_3$  addition was found to generally decrease enzyme  
546 activities for *C. vulgaris*. Additionally, both forms of N addition led to  
547 reductions in chitinase activity, which is similar to the findings of  
548 Bragazza et al. (2006), who suggested that this indicates an alleviation  
549 of N limitation.

550  
551 In *C. vulgaris* roots  $\beta$ -xylosidase,  $\beta$ -glucuronidase, and  $\beta$ -glucosidase  
552 activities under  $\text{NH}_4\text{Cl}$  addition were comparable to or higher than their  
553 activities in controls and  $\text{NaNO}_3$ +PK treatments, suggesting that  $\text{NH}_4\text{Cl}$   
554 promotes overall decomposition activity, as these enzymes primarily  
555 degrade plant cell wall components (Dunn et al., 2014). This may reflect  
556 a reduction in N limitation for the *C. vulgaris* mycorrhizal symbiont  
557 identified as *Hyaloscypha hepaticicola*, as experimental *in vitro* data has  
558 shown that this species may preferentially utilize  $\text{NH}_4^+$  as a source of N  
559 compared to organic sources (Cairney et al., 2000). Additionally, this is  
560 supported by the colonization morphotype data showing that  $\text{NH}_4\text{Cl}$   
561 addition induced the largest significant increase in intracellular ERM  
562 hyphae in *C. vulgaris*, implying increases in fungal biomass and

563 therefore higher decomposition potential. Furthermore, the sequence  
564 identification of *H. hepaticicola* in a *C. vulgaris* root sample from the  
565  $\text{NH}_4\text{Cl}$  treatment was linked to relatively higher enzymatic activities  
566 compared to other root samples in the same treatment. *Calluna vulgaris*  
567 roots hosting *Hyaloscypha* sp. displayed higher enzymatic activity under  
568 nutrient addition than in control conditions, indicating a response to  
569 increased N or NPK availability.

570

571 Our results demonstrate that it is necessary for studies of mycorrhizal  
572 fungi to include measurements of enzyme activities in natural  
573 conditions in order to more precisely estimate their roles in nutrient  
574 cycles. Though there is extensive research on the enzymatic activity of  
575 mycorrhizal fungi in sterile systems, few studies have measured the  
576 activity of mycorrhizal roots in their natural environment. While these *in*  
577 *vitro* enzyme activities of mycorrhizal fungi are often interpreted as  
578 their natural activities, work by Timonen and Sen (1998) showed that  
579 enzyme expression levels in *Pinus sylvestris* mycorrhizal fungi are locally  
580 regulated in the mycorrhizosphere, highlighting the variability in fungal  
581 enzyme expression which is not apparent from *in vitro* studies.

582

583 We observed that long-term nutrient addition resulted in a reduction in  
584 *C. vulgaris* abundance, potentially due to a reduction in competitive  
585 fitness, leaving an ecosystem gap that was rapidly occupied by other  
586 fast-growing species such as the non-mycorrhizal sedge *Eriophorum*

587 *vaginatum*, which is not reliant on symbionts for organic N uptake  
588 (Chapin et al., 1993). The suppressive nature of  $\text{NaNO}_3$  on *C. vulgaris*  
589 root enzymatic activities, compared to *E. tetralix*, suggests that *C.*  
590 *vulgaris* and its symbionts are more sensitive to  $\text{NaNO}_3$  and its effects  
591 on peat properties, such as pH. This is also reflected in the vegetation  
592 abundance data for  $\text{NaNO}_3$  where *C. vulgaris* abundance is reduced  
593 while *E. tetralix* abundance increases. This sensitivity to  $\text{NaNO}_3$  may put  
594 *C. vulgaris* at a competitive disadvantage to other ericaceous species  
595 during  $\text{NO}_3^-$  deposition.  
596  
597 Our findings of the cumulative effects of nutrient addition treatments at  
598 Whim Bog on abundances of peatland vegetation are similar to those  
599 detailed in Levy et al. (2019) which describes the decline of several plant  
600 species and the increase of *E. vaginatum* as the major effects over the  
601 entire timespan of the experimental site. The loss of *Sphagnum* may  
602 also directly reduce the ability of ericaceous species to uptake nutrients,  
603 as the upper moss layer is heavily inhabited by ericaceous roots,  
604 forming a thick layer which receives nutrients from litter and the  
605 atmosphere before it reaches the lower layers (Read et al., 2004). As the  
606 living *Sphagnum* layer is lost and forms bare, decaying peat, it collapses  
607 and becomes more submerged and anoxic, becoming an environment  
608 that ericaceous roots are less likely to inhabit. This loss of aerobic  
609 substrate for ericaceous species to inhabit and uptake nutrients from  
610 may be an underlying cause of the observed reduction in *C. vulgaris*

611 abundance. The loss of the moss layer may also lead to the subsidence  
612 of peat, as has been observed by Juutinen et al. (2018) to be a result of  
613 nutrient addition at another long-term nutrient addition experiment,  
614 Mer Bleue Bog, located in Ontario, Canada. This subsidence was  
615 indicated by our water table depth measurements, as they were made  
616 relative to the moss surface and the treatments with the highest water  
617 table values,  $\text{NaNO}_3$  and  $\text{NaNO}_3+\text{PK}$ , also showed the largest reductions  
618 in moss abundance. Furthermore, the effects of the treatments on peat  
619 pH should not be overlooked as a significant source of variation. Long-  
620 term alteration of pH is likely directly linked to the observed differences  
621 in fungal colonization patterns as well as root enzymatic activity.  
622  
623 Loss of ericaceous vegetation and *Sphagnum* are key examples of the  
624 risks posed by anthropogenic N and P deposition. Current research has  
625 lacked a clear picture of how ericaceous root associated fungi, ERM and  
626 DSE, are involved in these processes. Our findings indicate that altered  
627 nutrient limitations shift root associated fungal diversity and  
628 morphology, with direct effects on enzyme activity and thereby  
629 decomposition potential. The losses of *C. vulgaris* and *Sphagnum* due to  
630 nutrient addition and the subsequent reduction in DSE colonization  
631 rates may have additional consequences. Dark septate endophytes are  
632 by nature heavily melanized and may contribute a significant source of  
633 recalcitrant C (Fernandez and Koide, 2014). Subsequently, the potential  
634 loss of recalcitrant fungal biomass may lead to lower peatland C

sequestration. As suggested by Averill et al. (2014) and Orwin et al. (2011), ERM symbionts may be responsible for limiting the decomposition potential of free-living saprotrophs and the microbial community by increasing N and P limitation in soil. Addition of inorganic N and P may upset this limitation, leading to increased decomposition which releases C limitation for the more efficient saprotrophs, which in turn outcompete the mycorrhizal fungi, thereby limiting ericaceous nutrient access. Further research is necessary to determine the potential long-term risks of N and P deposition and the role of peatland mycorrhizal fungi in C sequestration.

645

#### 4.1 Conclusions

The total frequency of fungal colonization at Whim Bog, Scotland, did not decrease under nutrient addition treatments but instead tended to increase by ca. 10% in both *Calluna vulgaris* and *Erica tetralix*, refuting our hypothesis (1) which expected a reduction in fungal colonization rates. The considerable increase in ERM hyphal frequency (ca. 20-30%) in both host species was accompanied by a significant suppression of DSE hyphal frequency (ca. 20-30%) under nutrient addition, indicating a strong treatment effect on the root symbiont community. The altered fungal morphotype frequencies and identified fungal species agree with our hypothesis (2) of mycorrhizal diversity indicating nutrient addition effects and suggests that altered nutrient availability is a selective pressure upon the root associated fungal community. The enzymatic

659 activities of both ericaceous shrub roots and their associated fungi  
660 strongly support our hypothesis (3) of mycorrhizal root enzyme  
661 activities reflecting nutrient limitations.

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#### 677 **Acknowledgements**

678 We thank Sirpa Tiikkainen, Minna Oksanen, and Tuija Hytönen for help  
679 with the Direct PCR analyses. The study was funded by the Academy of  
680 Finland (projects 286731, 293365, 319262 to TL). The visits to Whim Bog  
681 were also supported by an eLTER H2020 Transnational Access grant to  
682 TL.



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923 **Figure Captions**

924 Figure 1. Total observed ericoid mycorrhizal (ERM) and dark septate  
 925 endophyte (DSE) occurrence in *C. vulgaris* and *E. tetralix* roots as  
 926 determined by light microscopy and the magnified intersections  
 927 method. Error bars indicate  $\pm 1$  standard deviation,  $n=3$ .

928  
 929 Figure 2. Microscopically observed frequencies of potential ericoid  
 930 mycorrhizal morphotypes (ERM1-4) in *C. vulgaris* and *E. tetralix* roots.  
 931 Error bars indicate  $\pm 1$  standard deviation,  $n=3$ . Different superscript  
 932 letters indicate significant differences ( $P<0.05$ ) compared with the other  
 933 treatments.

934  
 935 Figure 3. Microscopically observed frequencies of dark septate  
 936 endophyte morphotypes (DSE1-4) in *C. vulgaris* and *E. tetralix* roots.  
 937 Error bars indicate  $\pm 1$  standard deviation,  $n=3$ . Different superscript  
 938 letters indicate significant differences ( $P<0.05$ ) compared with the other  
 939 treatments.

940  
 941 Figure 4. Correlation analysis of *Calluna vulgaris* root enzyme activities  
 942 (Leu: leucine aminopeptidase, Glr:  $\beta$ -glucuronidase, Xyl:  $\beta$ -xylosidase,  
 943 Cell: cellobiohydrolase, Glc:  $\beta$ -glucosidase, Nag: N-  
 944 acetylglucosaminidase, Pho: acid phosphatase), root associated fungal  
 945 morphotype categories ERM1-4 and DSE1-4 (Table 1), surface peat (0-  
 946 20cm) pH, plot water table depth (WT), *C. vulgaris* abundance (Cv), *Erica*

947 *tetralix* abundance (Et), *Eriophorum vaginatum* abundance (Et), other  
 948 vascular spp. abundance (OV), *Sphagnum* abundance (Sph), and other  
 949 moss spp. abundance (OM). Statistically significant and indicative  
 950 correlation values are indicated by asterisks (\*\* for  $p<0.01$ , \* for  $p<0.05$ ,  
 951 and + for  $p<0.1$ ) values, with a greyscale ranging from lighter shades  
 952 indicating positive correlation coefficients to darker indicating negative  
 953 correlation coefficients,  $n=15$ .

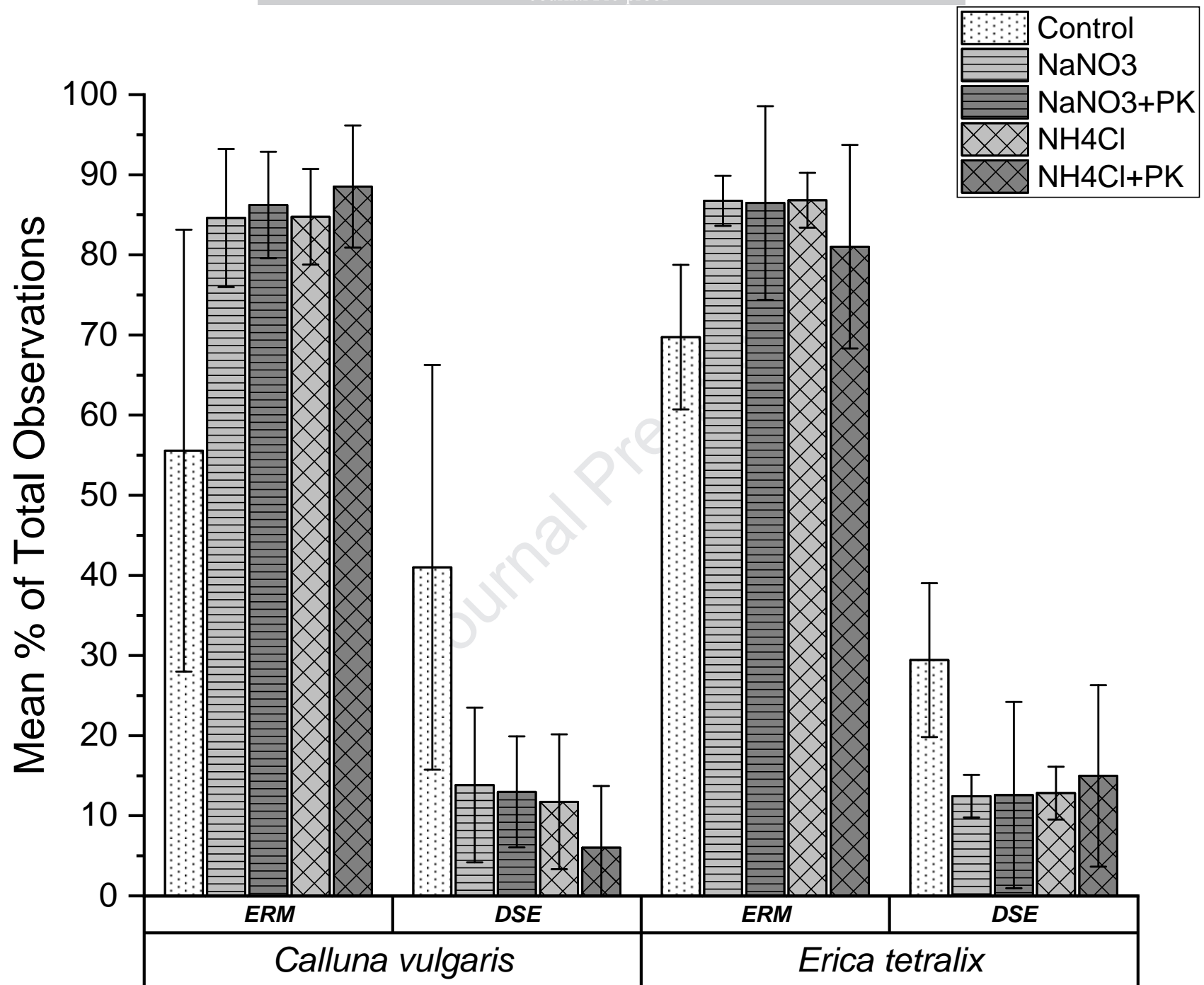
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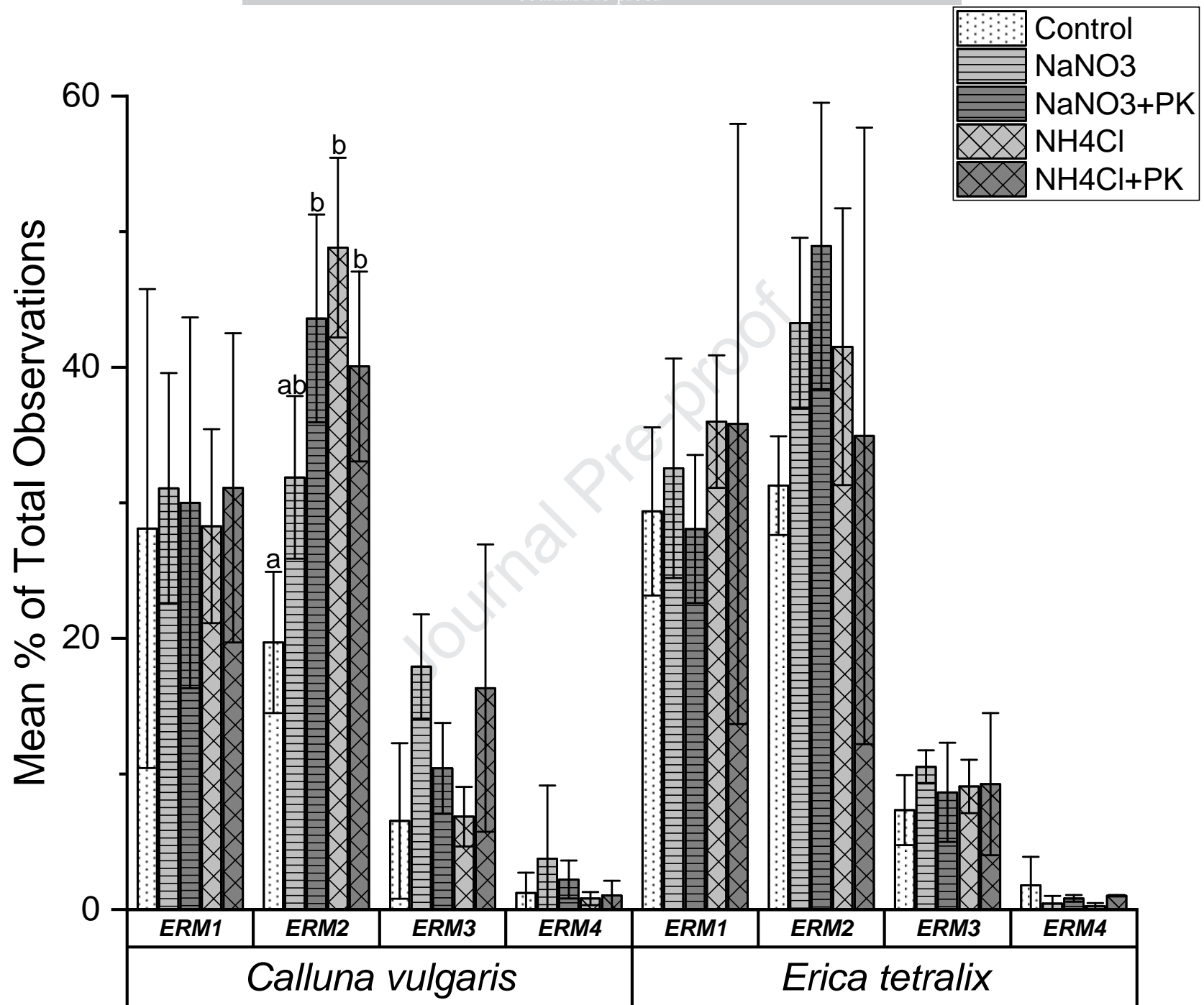
955 Figure 5. Correlation analysis of *Erica tetralix* root enzyme activities  
 956 (Leu: leucine aminopeptidase, Glr:  $\beta$ -glucuronidase, Xyl:  $\beta$ -xylosidase,  
 957 Cell: cellobiohydrolase, Glc:  $\beta$ -glucosidase, Nag: N-  
 958 acetylglucosaminidase, Pho: acid phosphatase), root associated fungal  
 959 morphotype categories ERM1-4 and DSE1-4 (Table 1), surface peat (0-  
 960 20cm) pH, plot water table depth (WT), *Calluna vulgaris* abundance  
 961 (Cv), *E. tetralix* abundance (Et), *Eriophorum vaginatum* abundance (Et),  
 962 other vascular spp. abundance (OV), *Sphagnum* abundance (Sph), and  
 963 other moss spp. abundance (OM). Statistically significant and indicative  
 964 correlation values are indicated by asterisks (\*\* for  $p<0.01$ , \* for  $p<0.05$ ,  
 965 and + for  $p<0.1$ ) values, with a greyscale ranging from lighter shades  
 966 indicating positive correlation coefficients to darker indicating negative  
 967 correlation coefficients,  $n=15$ .

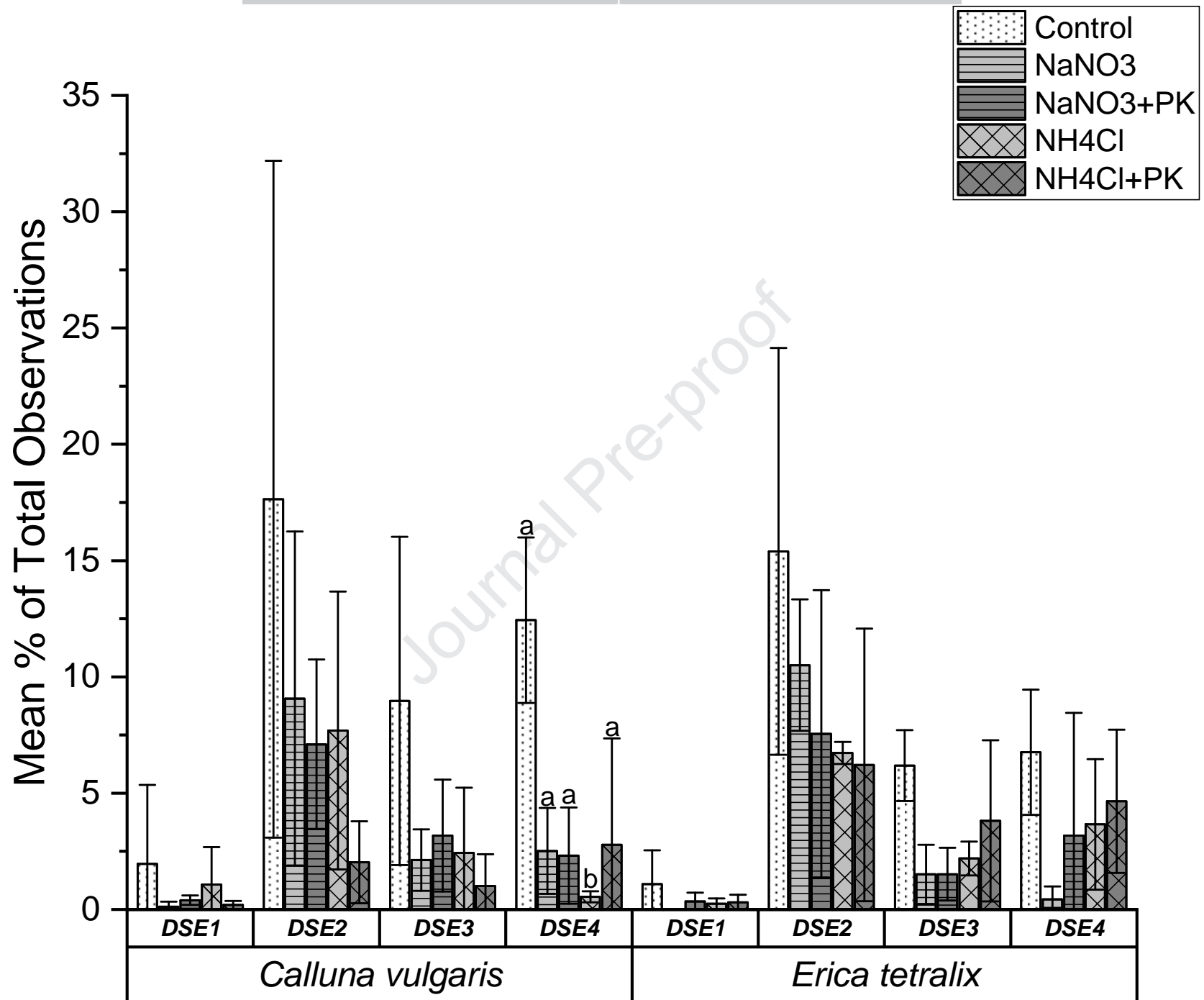
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969 Figure 6. *C. vulgaris* and *E. tetralix* root surface enzyme activities  
 970 according to treatment means (Leu: leucine aminopeptidase, Xyl:  $\beta$ -

971 xylosidase, Glr:  $\beta$ -glucuronidase, Cell: cellobiohydrolase). Error bars  
 972 indicate  $\pm 1$  standard deviation,  $n=3$ .  
 973  
 974 Figure 7. *C. vulgaris* and *E. tetralix* root surface enzyme activities  
 975 according to treatment means (Nag: N-acetylglucosaminidase, Glc:  $\beta$ -  
 976 glucosidase, Pho: acid phosphatase). Error bars indicate  $\pm 1$  standard  
 977 deviation,  $n=3$ . Letters above error bars indicate statistically significant  
 978 differences.  
 979  
 980 Figure 8. Ericaceous root enzyme activities (Leu: leucine  
 981 aminopeptidase, Glr:  $\beta$ -glucuronidase, Xyl:  $\beta$ -xylosidase, Cell:  
 982 cellobiohydrolase, Glc:  $\beta$ -glucosidase, Nag: N-acetylglucosaminidase,  
 983 Pho: acid phosphatase) for individual root pieces with identified fungal  
 984 sequences, as well as the mean activities for root samples without  
 985 sequencing (Black X Symbol,  $20 < n < 25$ , per treatment). Error bars  
 986 indicate  $\pm 1$  standard deviation of the samples without sequencing. Not  
 987 shown; *Acrodontium crateriforme*: Xyl=0.61, Nag=7.5, (*C. vulgaris*,  
 988  $\text{NH}_4\text{Cl}+\text{PK}$ ). *Pseudogymnoascus* sp.: Cell=0.87, (*C. vulgaris*,  $\text{NaNO}_3+\text{PK}$ ).

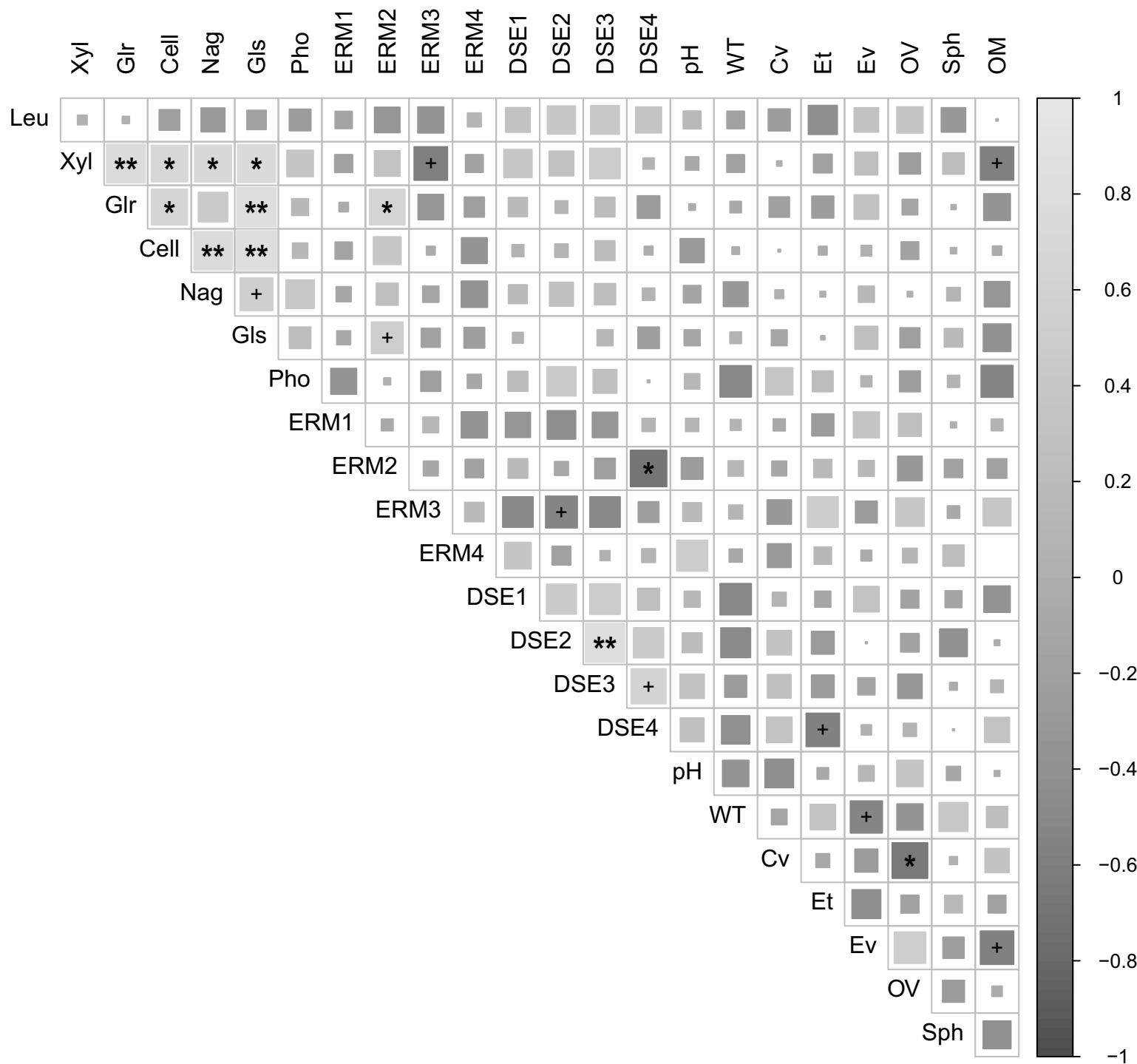




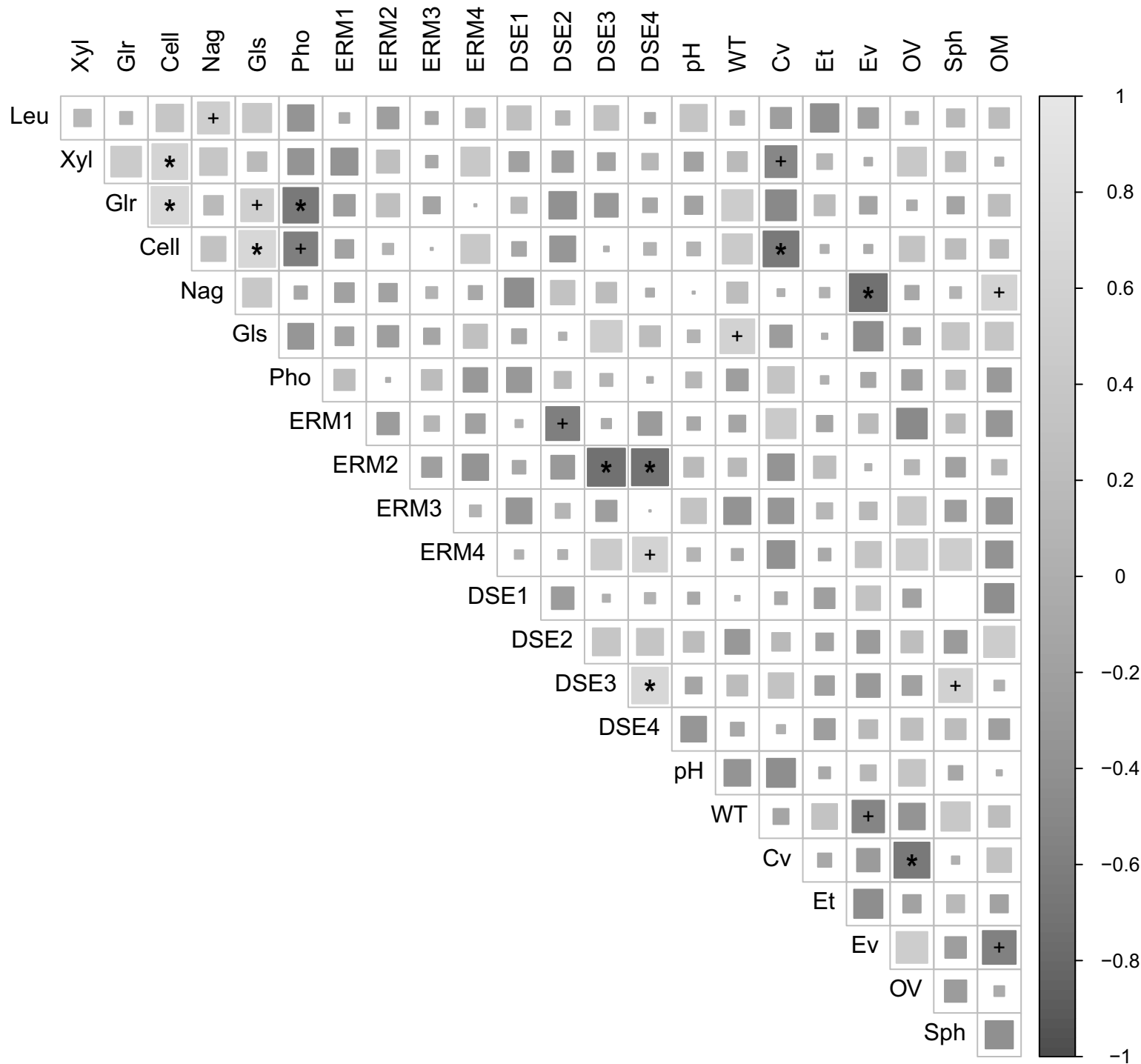


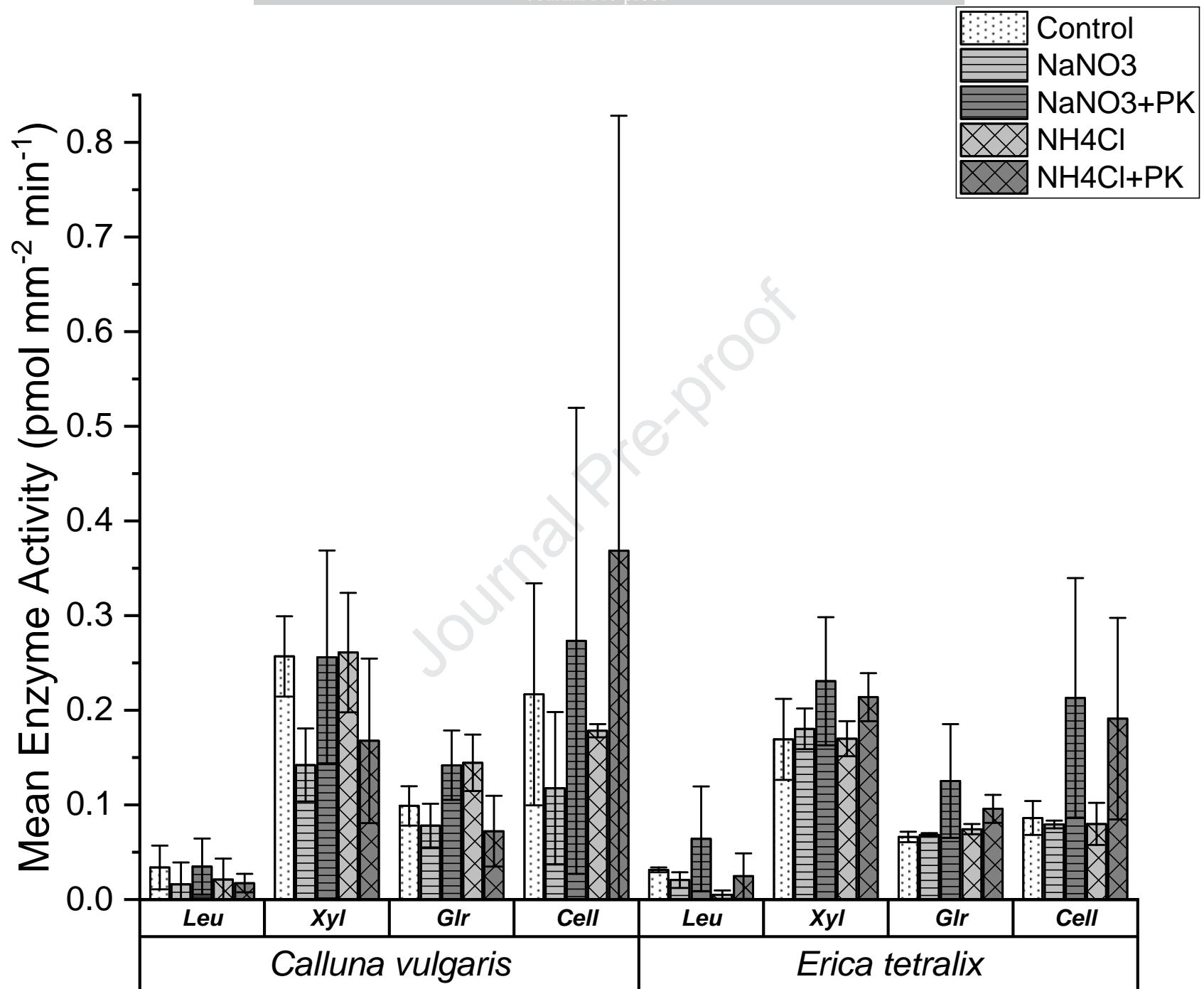


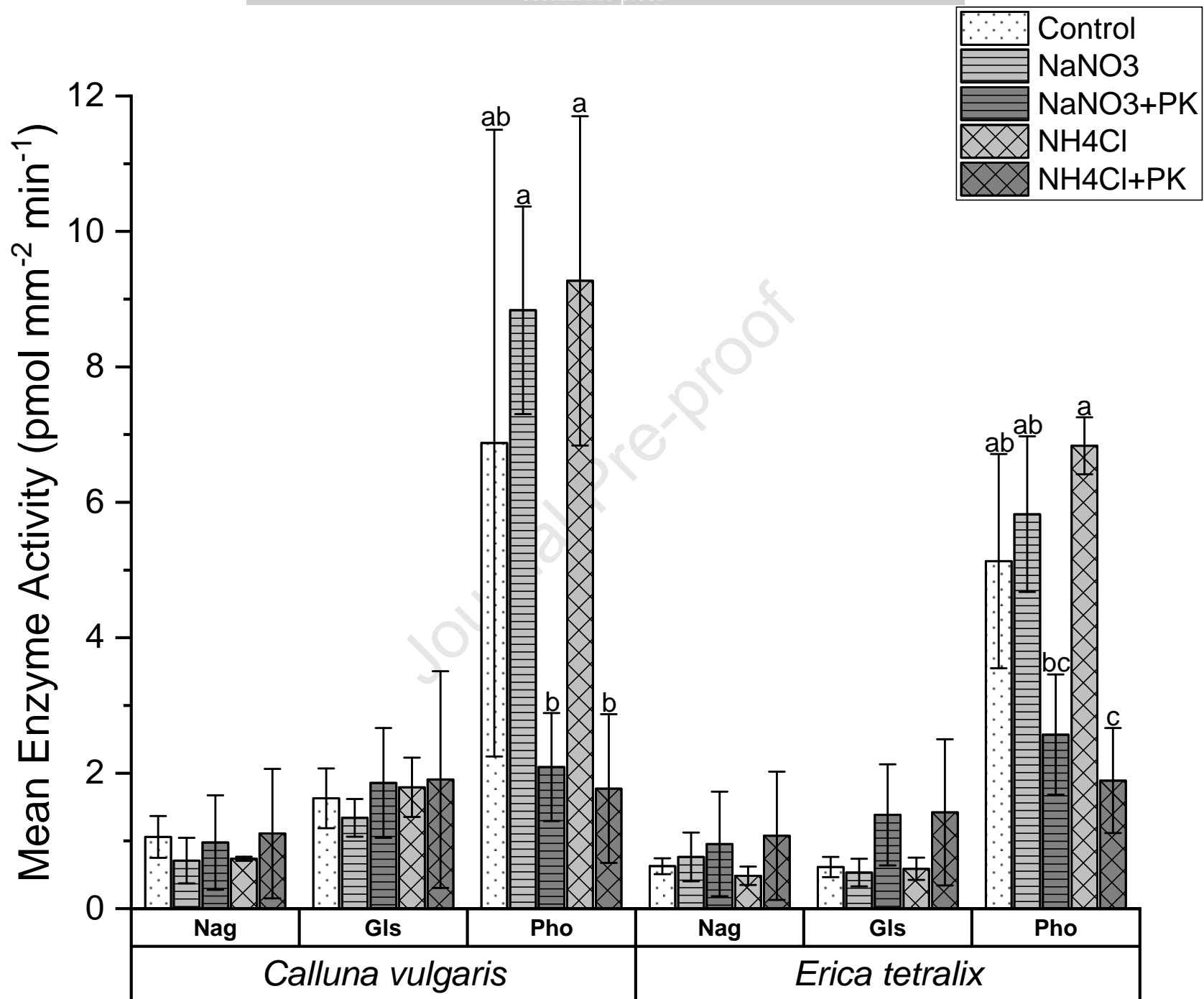
## ***Calluna vulgaris* Correlations**

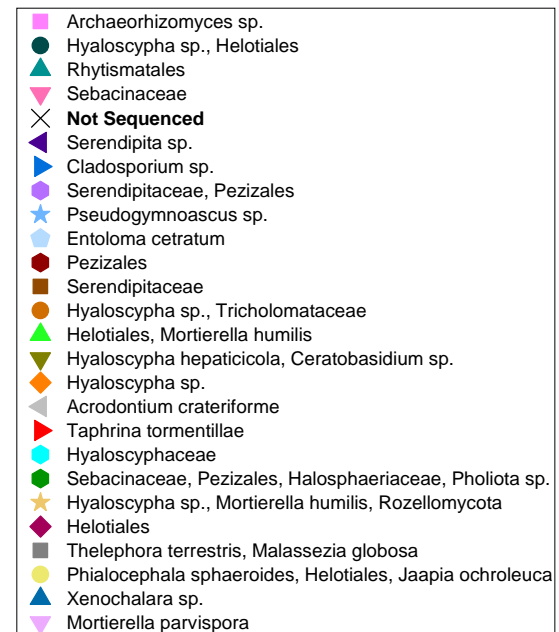
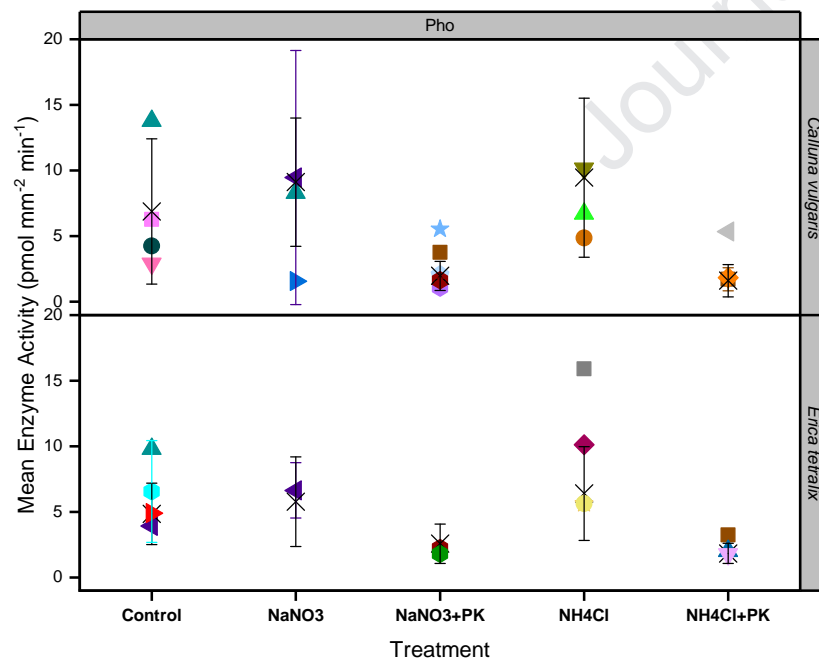
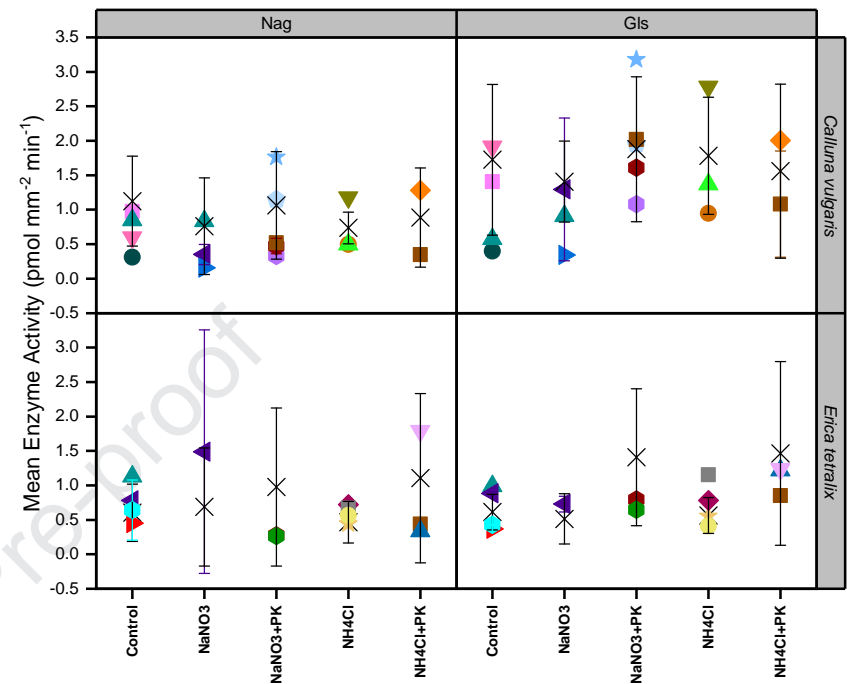
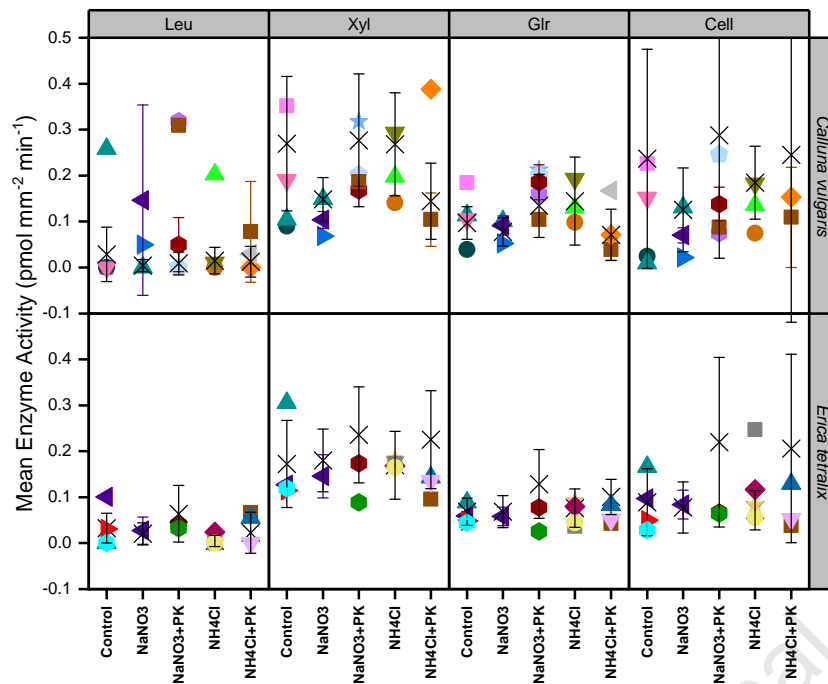


## *Erica tetralix* Correlations









- Long-term effects of N/NP on ericaceous root fungal colonization & enzymatic activity
- ERM intracellular colonization unchanged, ERM hyphae increased & DSE hyphae decreased
- Loss of recalcitrant fungal biomass in melanized DSE may reduce peatland C sink
- Nitrate reduced *C. vulgaris* root enzyme activity, suggests host/symbiont sensitivity

**Declaration of interests**

☒ The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

☐ The authors declare the following financial interests/personal relationships which may be considered as potential competing interests: